

**National Institute of Mental Health**  
*ClinicalTrials.gov*  
**Reference Guide for NIMH Researchers**  
**Revised April 2008**

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## Background

### What is *ClinicalTrials.gov*?

*ClinicalTrials.gov* is an online registry (<http://clinicaltrials.gov>), developed and maintained by the National Library of Medicine (NLM), that makes basic information on human research studies available to the public. This free registry currently contains descriptions of more than 50,000 federally and privately supported trials from 154 countries. Each record reflects a single unique protocol (e.g., a single record for a multi-site study), and includes a protocol summary describing the purpose of the study, recruiting status, criteria for patient participation, all location(s) of the trial, and a contact for additional information. All trials, regardless of study design, sponsor, and intervention types (e.g., drug, device, observational, behavioral), are accepted by *ClinicalTrials.gov*.

It is important for NIMH researchers to register their NIMH-supported clinical trials for several reasons. Registration is federally mandated for certain types of studies (see FDA Amendments Act of 2007, <http://prsinfo.clinicaltrials.gov/fdaaa.html>; NIH Guidance, <http://grants.nih.gov/grants/guide/notice-files/NOT-OD-08-023.html> and [http://grants.nih.gov/grants/policy/hs/faqs\\_aps\\_clinical\\_trials.htm](http://grants.nih.gov/grants/policy/hs/faqs_aps_clinical_trials.htm)). Additionally, scientific journals represented by the International Committee of Medical Journal Editors (ICMJE) require studies to be registered as a prerequisite to publication of the trial results (N Engl J Med 2004; 351:1250-1; <http://www.icmje.org>). Finally, studies registered with *ClinicalTrials.gov* have the potential to be seen by a large audience of referring clinicians and potential trial participants and their family members. The registry enables visitors to search for clinical trials by location, thus facilitating instant results for regional queries.

#### Additional Resources:

- Information for NIMH investigators conducting studies: <http://www.nimh.nih.gov/studies/researchers.cfm>
- *ClinicalTrials.gov* FAQs: <http://www.nlm.nih.gov/services/faqctgov.html>
- About *ClinicalTrials.gov*: <http://www.clinicaltrials.gov/ct2/info/about>
- Center for Drug Evaluation and Research (CDER) Guidelines: <http://www.fda.gov/cder/guidance/4856fnl.htm>

# Registering and Maintaining Your NIMH-Sponsored *ClinicalTrials.gov* Record

## Getting Started

Once a study has been funded by NIMH, it is eligible for inclusion in *ClinicalTrials.gov*.

Study records are registered and maintained through the Protocol Registration System (PRS), a web-based tool developed for managing clinical trial information submissions. Records submitted through the PRS (<http://register.clinicaltrials.gov>) will be made public, after review and approval, in the NLM's *ClinicalTrials.gov* public website (<http://clinicaltrials.gov>).

The NIMH *ClinicalTrials.gov* Records Coordinator can help facilitate the registration and/or maintenance of a clinical trial in *ClinicalTrials.gov*. If the PI or study designee is a first-time user of the PRS, the PI will be contacted by the Records Coordinator via email, with information about content requirements, instructions about how to register, and a User Name and Password to be used to log into the PRS.

PRS users enter their own information about their clinical trials. Users should ensure that the information is correct, easily understood by members of the public, and updated in a timely manner. The *ClinicalTrials.gov* team maintains PRS and the *ClinicalTrials.gov* website and may make minor modifications to trial records for clarity.

After the PI or designated study official registers with PRS, he/she may open his/her record and, following the PRS Data Element Definitions, provide information for the mandatory fields. See **Appendix 1** for additional information on these fields.

## Completing the Study Record

Once the clinical trial record fields are completed and the updater clicks [Complete] at the top of the screen, PRS automatically notifies the NIMH *ClinicalTrials.gov* Records Coordinator, who reviews the record for completeness. The record is then sent to the NIMH *ClinicalTrials.gov* Administrator for approval. After the record is released by NIMH, the NLM publishes it on the *ClinicalTrials.gov* website. See **Appendix 2** for an overview of the process for registering trials.

## Is it necessary to seek IRB approval for information listed in the *ClinicalTrials.gov* record?

The information submitted for the *ClinicalTrials.gov* record that describes and summarizes the clinical trial does not require IRB approval.

Please refer to section L of the FDA's "Guidance for Industry Program on Clinical Trials for Serious or Life-Threatening Diseases and Conditions," which states:

***“Is Institutional Review Board preapproval of the protocol listing required?”***

*No. Section 113 of the Modernization Act does not require prior IRB approval when submitting this information to the Clinical Trials Data Bank. Current FDA guidance*

recommends that IRB review of listings need not occur when, as here, the system format limits the information provided to basic information, such as title, purpose of the study, protocol summary, basic eligibility criteria, study site locations, and how to contact the site for further information.” <http://www.fda.gov/cder/guidance/4856fnl.htm>

## **Once an NIMH-sponsored study is registered in *ClinicalTrials.gov*, who is responsible for maintaining and updating the information about that study?**

The PI or a designated study representative is the most appropriate person for updating and maintaining data about the study. The study officials are responsible for the completeness and updating of the record once it becomes part of the system. They serve as a point of contact for the *ClinicalTrials.gov* team and resolve questions associated with the information that is provided. When protocol or any other information included in the *ClinicalTrials.gov* record changes, it is the responsibility of the PI or study representative to update the record to reflect the change.

NOTE: Each *ClinicalTrials.gov* record has one “owner”. This owner is the only study individual responsible for and authorized to make changes to the record. If the “owner” of a study record changes, the “owner” must contact the Records Coordinator or the NIMH Administrator to change the record ownership. The record owner should designate the new person who will be responsible for maintaining the study record, whether this person is the PI, study coordinator, or central contact of the study. Ownership of the account will then be transferred to the designated person.

## **Does study information need to be updated after a trial has closed?**

Yes, a *ClinicalTrials.gov* record is permanent and will always be displayed in *ClinicalTrials.gov*, regardless of the trial’s status. However, after a trial’s status has been changed to “completed,” study contact information will no longer be displayed. It is the investigator’s responsibility to make sure the *ClinicalTrials.gov* record is accurate, even if the study has closed. This includes adding journal citations of the study’s results. For an example, see “Publications that report results of this study” at the bottom of the CATIE schizophrenia trial record: <http://www.clinicaltrials.gov/ct/show/NCT00014001>.

If the journal is indexed in PubMed, an automatic link to the publication will appear on your study’s *ClinicalTrials.gov* page after you have added the citation’s PubMed Identifier (PMID) to your study record. If the journal in which your results are published is not indexed in PubMed, you may still add the citation, but it will not produce a link to your publication.

Reminders about updating records are periodically sent to record owners. For trials listed as “Not Yet Recruiting,” “Recruiting,” and “Suspended,” reminders are sent twice a year. Reminders are sent to records with the recruiting status “Active, Not Recruiting” once a year.

## Appendix 1: Selected PRS Data Element Definitions with NIMH-Specific Examples

The following information has been adapted from the NLM's online PRS User's Guide and is available in full in the *ClinicalTrials.gov* PRS Guidelines (<http://prsinfo.clinicaltrials.gov/definitions.html>).

### User Definitions

#### PRS Administrators

- NIMH *ClinicalTrials.gov* PRS Records Coordinator: responsible for working with Principal Investigator (PI) to create, manage, and maintain *ClinicalTrials.gov* records (Meghan Bishop: [update@clinicaltrials.gov](mailto:update@clinicaltrials.gov))
- NIMH *ClinicalTrials.gov* Administrator: responsible for reviewing, approving, and releasing records for publication on *ClinicalTrials.gov* (Jean Baum: [jbaum@mail.nih.gov](mailto:jbaum@mail.nih.gov) and Christine Ulbricht: [uchristi@mail.nih.gov](mailto:uchristi@mail.nih.gov))

#### PRS Users

- Owner: responsible for creating and updating the record as necessary (the PI or designated primary updater of the record)
- Updater: the person who last updated the record (usually the owner, the Records Coordinator, or the Administrator)

### User Responsibilities

PRS users provide and maintain information about their clinical trials by entering information into PRS and ensuring that the information is correct, easy to understand, and updated in a timely manner. Through PRS, a user may:

- Enter information regarding clinical trials
- Modify a record
- View a record
- Change a password
- Preview a record as it will appear on *ClinicalTrials.gov*
- Complete and submit the trial data for approval

### Special Cases

#### Multi-site Studies

For trials being conducted at multiple study sites under different PIs, only one record should be created in *ClinicalTrials.gov*. In order to avoid the duplication of records, a Central Contact (a PI or another study official) needs to be designated to take primary responsibility for entering information from all of the study sites. All PIs involved in a multi-site trial will be given the contact information for the Central Contact. PIs will be responsible for sending the Central Contact any updates related to their respective sites.

## **Continuation Awards**

For studies that are recipients of awards covering multiple years of work, a record may already have been created for a previous year. Please check in *ClinicalTrials.gov* to ensure that there are no duplicates before you create a record. You can search by entering any of the following into the search field on the *ClinicalTrials.gov* home page: title of the study, the grant number, a study official's name, or any other information that is unique to your study.

## **I. Titles and Background Information**

### **Organization's Unique Protocol ID**

Definition: Unique identification assigned to the protocol by the sponsoring organization, which will be the NIMH grant, contract, or cooperative agreement number. Multiple studies conducted under the same grant or contract must each have a unique number. The grant/contract number should be formatted in the following manner:

**For a grant funding only one study: R01 MH12345**

**For a contract: N01 MH90003**

For a multi-site study, enter the grant/contract number for the main study site as the Unique Protocol ID. Grant/contract numbers from other sites should be listed as secondary IDs.

### **Secondary IDs**

Definition: Other identification numbers assigned to the protocol, including any applicable NIH grant numbers.

**Examples: R01 MH61686-05; R01 MH059542; R01 MH059552; R01 MH075131; R01 MH059541; R01 MH060912; DNBBS 7G-GRR\***

\*Please note: NIMH will add the applicable division name and program class code (PCC) to the record.

### **Brief Title**

Definition: Protocol title written in plain language for the general public.

### **Official Title**

Definition: Official name of the protocol provided by the principal investigator or sponsor.

#### **Examples:**

- Brief Title:** Stimulant Versus Nonstimulant Medication for Attention Deficit Hyperactivity Disorder in Children  
**Official Title:** Measuring and Predicting Response to Atomoxetine and Methylphenidate
- Brief Title:** Behavioral Treatments for Acute Stress Disorder In Firefighters  
**Official Title:** Developing Group Treatments for Acute Stress Disorder
- Brief Title:** Characteristics of Sleep Patterns in Young Adults With and Without Insomnia  
**Official Title:** Psychobiology and Treatment Response in Primary Insomnia
- Brief Title:** Treatment of Mania Symptoms With Drug Therapy  
**Official Title:** Divalproex Extended Release and Placebo, Lithium, or Quetiapine for Mania

5. **Brief Title:** Effectiveness of Behavioral Treatments for Obesity and Major Depression in Women

**Official Title:** Treating Co-Morbid Obesity and Major Depressive Disorder

### Study Type

Definition: Nature of the investigation. Select one.

- **Interventional:** studies in human beings in which individuals are assigned to receive specific interventions. Subjects may receive diagnostic, therapeutic, or other types of interventions. The assignment of the intervention may or may not be random. The individuals are then followed and biomedical and/or health outcomes are assessed.
- **Observational:** studies in human beings in which biomedical and/or health outcomes are assessed in a predefined group of individuals. Subjects in the study may receive diagnostic, therapeutic, or other interventions, but the investigator does not assign specific interventions to the subjects of the study.
- **Expanded Access:** records describing the procedure for obtaining an experimental drug or device for patients who are not adequately treated by existing therapy, who do not meet the eligibility criteria for enrollment, or who are otherwise unable to participate in a controlled clinical study. Expanded Access records are used to register all types of non-protocol access to experimental treatments, including protocol exception, single-patient IND, treatment IND, compassionate use, emergency use, continued access, and parallel track.

## II. Sponsors

### Sponsor

Definition: Name of organization that is funding the clinical investigation.

**Note:** **National Institute of Mental Health** should be listed as the sponsor for all NIMH studies registered in PRS.

### Responsible Party

Definition: As defined in **US Public Law 110-85**, Title VIII, Section 801, the term "responsible party," with respect to a clinical trial, means

1. the sponsor of the clinical trial (as defined in 21 CFR 50.3) or
2. the principal investigator of the clinical trial if designated by a sponsor, grantee, contractor, or awardee, so long as the principal investigator is responsible for conducting the trial, has access to and control over the data from the clinical trial, has the right to publish the results of the trial, and has the ability to meet all of the requirements for the submission of clinical trial information.

Provide the following information for the designated responsible party:

1. Name/Official Title - for either the principal investigator or sponsor contact
2. Organization - the sponsor or the principal investigator's organizational affiliation
3. Contact Information - telephone number and/or email address (*required for internal administrative use only; not revealed to public*)

**Example:** Name/Official Title: John Smith, PhD  
Organization: Mount Sinai School of Medicine  
Phone: 555-920-1552 Ext: Email: [john.smith@mssm.edu](mailto:john.smith@mssm.edu)

### III. Human Subjects Review

Submitted studies must have approval from a human subjects review board, such as an Institutional Review Board (IRB), ethics committee, or equivalent group that is responsible for reviewing and monitoring human subjects in this protocol.

Review board information is not required for trials associated with U.S. FDA IND or IDE applications.

**Review board information is required for internal administrative use and is not revealed to the public.** Oversight authority information is displayed on *ClinicalTrials.gov*. For more details on this, please refer to the PRS *ClinicalTrials.gov* User's Guide online Data Element Definitions in the PRS HELP function.

#### **Oversight Authorities**

Definition: The name of each national or international health organization with authority over the protocol. Use the following format for each authority:

Country: Organization Name

**Example:**

*United States:* Federal Government

### IV. Study Description

**Brief Summary** (known as Purpose in the published record)

Definition: Short description of the primary purpose of the protocol intended for the lay public.

**Example 1: A Behavioral Intervention**

This study will determine the effectiveness of a group-based behavioral program for weight reduction in overweight and obese people with schizophrenia.

**Example 2: A Drug Intervention**

This study will compare two different antidepressant treatment regimens to determine which is more effective in reducing symptoms of bipolar depression.

**Example 3: An Observational Study**

This study will examine brain responses associated with reinforcement and reward tasks in individuals with major depressive disorder (MDD).

#### **Detailed Description**

Definition: Extended description of the protocol, including information not already contained in other fields. Generally, the description should contain two paragraphs. The first paragraph should include the rationale for the study, and the second paragraph should outline the methodology and duration of the study.

**Example 1: A Behavioral Intervention**

Somatization disorder is a chronic psychological condition that causes numerous physical complaints for which no underlying physical problem can be identified. The disorder often lasts for several years and results in substantial functional impairment. The physical complaints most frequently involve chronic pain and problems with the digestive, nervous, and reproductive systems. Neither pharmacological nor psychosocial treatments for this disorder have been successful in suppressing symptoms. Cognitive behavioral therapy (CBT) is a treatment that focuses on maladaptive patterns of thinking and the beliefs that underlie

such thinking. This study will examine the long-term effects of CBT on the physical symptoms, functioning, and health care utilization of people with somatization disorder.

Participants in this open label study will be randomly assigned to receive either CBT supplemented with augmented standard medical care (ASMC) as indicated by a psychiatric consultation letter or ASMC alone. Participants assigned to CBT plus ASMC will receive CBT for 10 weeks. Somatic symptomatology, functional impairment, and health care costs will be assessed at study visits at baseline and Months 3, 9, and 15. The visits at Months 9 and 15 will assess specifically the long-term efficacy of the treatment.

### ***Example 2: A Drug Intervention***

Generalized social anxiety disorder (GSAD) is one of the most common psychiatric disorders, and often causes significant distress and dysfunction in affected individuals. Although currently available treatments for GSAD are effective, most individuals have residual symptoms after initial psychosocial or psychopharmacologic intervention. Further treatment is necessary for such individuals, but sufficient research has not been done to guide clinicians on what the safest and most effective next step may be. This study will compare the effectiveness of either combining clonazepam or placebo with sertraline or completely switching to venlafaxine in treating GSAD in individuals who have not responded to treatment with sertraline. This study will also examine predictors of treatment response, including factors such as age at disease onset, duration of illness, comorbidities, and genes that influence serotonin and catecholamine metabolism.

Participants in this double-blind study will first partake in an initial 10-week phase in which they will be treated with sertraline. Participants who do not respond to sertraline treatment will proceed to phase two of the study, in which they will be randomly assigned to one of three treatment groups. One group will receive both sertraline and clonazepam, another group will receive both sertraline and placebo, and the third group will receive only venlafaxine. All treatments will continue for 12 weeks. Sertraline and venlafaxine are both FDA-approved for the treatment of GSAD. Clonazepam is widely used for the treatment of anxiety, but is not FDA-approved for the treatment of GSAD. All participants will attend weekly study visits at Weeks 1, 2, 4, 6, 8, and 10. Participants who continue into phase two will attend weekly study visits at Weeks 11 – 14, 16, 18, 20, and 22. Symptom remission rates and post-treatment social phobia severity will be assessed at Week 20.

### ***Example 3: An Observational Study***

Bipolar disorder (BPD), also known as manic-depressive illness, is a disorder that causes frequent shifts in an individual's mood, energy, and ability to function. An individual with BPD may go through periods of mania, which are characterized by increased energy, irritability, and an excessively "high" euphoric mood. The manic periods are followed by periods of depression, which are characterized by decreased energy, feelings of hopelessness, and anxiety. BPD is a persistent and severe mental illness with a high suicide rate; it must be strictly managed through medication and therapy. Many BPD medications have been developed recently; however, there are still many individuals who do not respond well to medication treatment. Research has shown that the way individuals experience illness has an effect on their response to medication. The purpose of this study is to gain insight into how individuals with BPD perceive and respond to medication treatment. Factors such as gender, degree of social support, drug and alcohol usage, and attitudes toward medication will be evaluated to understand how they affect medication and treatment adherence.

This 6-month study will consist of 3 interviews. Each interview will last approximately 2 and ½ hours and will include numerous standardized psychological questionnaires. The

questionnaires will assess participants' attitudes toward BPD treatment; psychiatric illness severity, including symptoms of mania and depression; level of addiction to alcohol and drugs; availability of social support resources; and medication adherence.

## V. Key Dates

### **Record Verification Date**

Definition: Date the protocol information, including recruiting status, was last verified, whether changes were made or not.

### **Study Start Date**

Definition: Date that enrollment to the protocol begins.

### **Primary Completion Date**

Definition: As specified in US Public Law 110-85, Title VIII, Section 801, with respect to an applicable clinical trial, the date that the final subject was examined or received an intervention for the purposes of final collection of data for the primary outcome, whether the clinical trial concluded according to the prespecified protocol or was terminated. A "Type" menu is also included, with options Anticipated and Actual. For active studies, set Type to Anticipated and specify the expected completion date, updating the date as needed over the course of the study.

### **Study Completion Date**

Definition: Final date on which data was (or is expected to be) collected. Use the Type menu (Anticipated/Actual) as described above.

## VI. Outcome Measures

### **Primary Outcome Measure**

Definition: Specific key measurement(s) or observation(s) used to measure the effect of experimental variables in a study, or for observational studies, to describe patterns of diseases or traits or associations with exposures, risk factors, or treatment.

### **Time Frame**

Definition: Time point(s) at which outcome measure is assessed.

### **Safety Issue?**

Definition: Is this outcome measure assessing a safety issue? Select: Yes/No.

### **Secondary Outcome Measures**

Definition: Other key measures that will be used to evaluate the intervention(s) or, for observational studies, that are a focus of the study. Specify Outcome Measure, Time Frame, and Safety Issue (See above).

#### ***Example 1:***

Primary Outcome Measure:

Measure: Clinician Severity Rating on the Anxiety Disorders Interview Schedule for Children (ADIS C/P)

Time Frame: Measured at pretreatment, midtreatment, and Year 1 follow-up post-treatment

Safety Issue?: No

Secondary Outcome Measures:

Measure: Child Manifest Anxiety Scale for Children, Revised  
Time Frame: Measured at pretreatment and Year 1 follow-up post-treatment  
Safety Issue?: No

**Example 2:**

Primary Outcome Measure:  
Measure: Time to medication discontinuation  
Time Frame: Measured at Year 1  
Safety Issue?: No

Secondary Outcome Measures:

Measure: Psychiatric symptoms, hospitalization, and medication side effects  
Time Frame: Measured at Year 1  
Safety Issue?: Yes

**Example 3 (Observational Study):**

Primary Outcome Measure:  
Measure: Social and emotional development of infants who have mothers with depression and anxiety, during or soon after pregnancy  
Time Frame: Measured at Months 6 and 12 post-pregnancy  
Safety Issue?: No

## VII. Arms, Groups, and Interventions

For interventional studies specify the arms:

### Arm Number or Label

Definition: The number, letter, or name used to identify the arm.

*Examples:* A, 2, III

### Arm Type

Select one:

- Experimental
- Active Comparator
- Placebo Comparator
- Sham Comparator
- No Intervention
- Other

### Arm Description

Definition: Brief description of the arm.

**Example 1:**

Participants will receive cognitive behavioral therapy (Arm 1: Experimental)  
Participants will receive treatment as usual (Arm 2: Active Comparator)

**Example 2:**

Participants will receive treatment with sertraline (Arm I: Experimental)  
Participants will receive treatment with placebo (Arm II: Placebo Comparator)

For observational studies specify the predefined participant groups (cohorts) to be studied. Do not use this section to specify strata (Detailed Design can be used for that purpose, if desired).

**Group/Cohort Number or Label**

Definition: The number, letter, or name used to identify the group.

*Examples:* A, 2, III, Surgical, Observation

**Group/Cohort Description**

Definition: Explanation of the nature of the study group (e.g., those with a condition and those without a condition; those with an exposure and those without an exposure). Note that the overall study population should be described under Eligibility.

*Examples:* Participants with depression (Group 1)  
Healthy participants without depression (Group 2)

For all studies, and for expanded access records, specify the associated intervention(s).

**Intervention Type**

Select one per intervention:

- Drug (including placebo)
- Device (including sham)
- Biological/Vaccine
- Procedure/Surgery
- Radiation
- Behavioral (e.g., psychotherapy, lifestyle counseling)
- Genetic (including gene transfer, stem cell, and recombinant DNA)
- Dietary Supplement (e.g., vitamins, minerals)
- Other

**Intervention Name**

Definition: For drugs use generic name; for other types of interventions provide a brief descriptive name.

For investigational new drugs that do not yet have a generic name, a chemical name, company code, or serial number may be used on a temporary basis. As soon as the generic name has been established, update the associated protocol records accordingly.

For non-drug intervention types, provide an intervention name with sufficient detail so that it can be distinguished from other similar interventions.

*Examples:* Cognitive behavioral therapy (CBT) (Behavioral)  
Fluoxetine (Drug)  
Placebo (Drug)

**Intervention Description**

Definition: Cover key details of the intervention. Must be sufficiently detailed to distinguish between arms of a study (e.g., comparison of different dosages of drug) and/or among similar interventions (e.g., comparison of multiple implantable cardiac defibrillators). For example, interventions involving drugs may include dosage form, dosage, frequency, and duration.

*Example 1:* Dosage ranging from 50 mg to 200 mg once a day for 12 weeks

**Example 2:** Participants will receive 16 interpersonal therapy (IPT) sessions, which will address adjustment to pregnancy, concerns about interpersonal relationships, and parenting issues.

### **Arms/Groups**

Definition: If arms or groups have been specified for the protocol, select the ones for which the intervention is to be administered. For interventional studies with arms specified, all arms must have at least one intervention (unless arm type is "No Intervention") and each intervention must be assigned to at least one arm. For observational studies with groups specified, each intervention (if any) must be assigned to at least one group.

### **Other Names**

Definition: List other names used to identify the intervention, past or present (e.g., brand name for a drug). These names will be used to improve search results in *ClinicalTrials.gov*.

### **Example 1 - Arms, Groups, and Interventions (Interventional Study):**

Arms:

- 1: Experimental  
Participants will receive sertraline and cognitive behavioral therapy
- 2: Active Comparator  
Participants will receive placebo and cognitive behavioral therapy

Interventions:

Drug: Sertraline

Sertraline will be administered in standard dosing or slow titration. Treatment with sertraline will last 18 weeks.

Arms: 1

Other Names: Zoloft

Drug: Placebo

The placebo will be administered in the same manner as sertraline. Treatment with placebo will last 18 weeks.

Arms: 2

Behavioral: Cognitive behavioral therapy (CBT)

CBT treatment will begin at Week 4 of antidepressant treatment. CBT will include education, training, and identifying repetitive behaviors of participants. Participants will learn how to respond to repetitive behaviors in a positive manner.

Arms: 1, 2

### **Example 2 - Arms, Groups, and Interventions (Observational Study):**

Groups/Cohorts:

- 1: Smokers with schizophrenia
- 2: Smokers with bipolar disorder
- 3: Smokers without any mental illness

Interventions:

Device: CReSSmicro handheld topography device

The CReSSmicro device represents the state-of-the-art technology for measurements of ambulatory puff topography taken in the smoker's natural environment. Although all topography measurements are limited, at least to some degree, by the artificial act of smoking while using a device, or smoking through a mouthpiece, this small, lightweight, and portable device is easy to use outside of the laboratory setting to capture more naturalistic smoking behavior and allows for less intrusion from the research team and research environment.

Groups: 1, 2, and 3

## VIII. Conditions and Keywords

### Conditions or Focus of Study

Definition: Primary disease or condition being studied, or focus of the study. When naming diseases or conditions, the NLM's Medical Subject Headings (MeSH) controlled vocabulary should be used when possible.

*Examples:* Anxiety Disorders  
Depression  
Bipolar Disorder  
Eating Disorders  
Generalized Anxiety Disorder

### Keywords

Definition: Words or phrases that best describe the protocol. Keywords help users find studies in the database. Use the NLM's Medical Subject Heading (MeSH) controlled vocabulary terms where appropriate. Be as specific and precise as possible. Avoid acronyms and abbreviations.

*Examples:* Major Depressive Disorder  
Psychosocial Intervention  
Anorexia

## VIII. Related Information

### References

Definition: Citations to publications related to the protocol: background and/or results. Provide either the unique PubMed Identifier (PMID) of an article or enter the full bibliographic citation.

### MEDLINE Identifier

Definition: Unique PubMed Identifier (PMID) for the citation.

*Example:* PMID: 12000823

### Citation

Definition: Bibliographic reference in NLM's MEDLINE format.

*Example:* Tolin, D.F., Diefenbach, G.J., Maltby, N., & Hannan, S. (2005). Stepped care for obsessive-compulsive disorder: A pilot study. *Cognitive and Behavioral Practice*, 12, 403-414.

### Results Reference

Definition: Indicate if the reference provided reports on results from this clinical research study.

NOTE: Currently active studies should not have results references. Results references should be added after study completion.

### **Links**

Definition: A Web site directly relevant to the protocol may be entered, if desired. Do not include sites whose primary goal is to advertise or sell commercial products or services. Links to educational, research, government, and other non-profit Web pages are acceptable. All submitted links are subject to review by *ClinicalTrials.gov*.

### **URL**

Definition: complete URL, including http://

### **Description**

Definition: Title or brief description of the linked page. If the page being linked is the protocol's home page on the sponsor's Web site, include the words "Click here for more information about this study:" and provide the name of the protocol.

### **Example:**

<http://www.catie.unc.edu/> (Click here for more information about this study)

### **Examples of NIMH study records in *ClinicalTrials.gov***

<http://www.clinicaltrials.gov/show/NCT00611975>

<http://www.clinicaltrials.gov/show/NCT00601393>

<http://www.clinicaltrials.gov/show/NCT00601653>

<http://www.clinicaltrials.gov/show/NCT00601965>

<http://www.clinicaltrials.gov/show/NCT00614068>

<http://www.clinicaltrials.gov/show/NCT00611806>

## Appendix 2: Process for Registering Trials in *ClinicalTrials.gov*

